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- **COMPOSITIONS AND METHODS FOR USE** (54)**IN THREE DIMENSIONAL MODEL** PRINTING
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#### **Related U.S. Application Data**

Continuation of application No. 12/963,577, filed on (63)Dec. 8, 2010, now Pat. No. 8,106,107, which is a continuation of application No. 12/342,210, filed on Dec. 23, 2008, now Pat. No. 7,851,122, which is a continuation of application No. 11/518,295, filed on Sep. 11, 2006, now Pat. No. 7,479,510, which is a continuation of application No. 11/098,690, filed on Apr. 5, 2005, now Pat. No. 7,183,335, which is a continuation of application No. 10/424,732, filed on Apr. 29, 2003, now abandoned, which is a continuation-in-part of application No. 09/803,108, filed on Mar. 12, 2001, now Pat. No. 6,569,373.

See application file for complete search history.

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| 8,106,107 B2* | 1/2012  | Napadensky 522/75       |

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#### ABSTRACT (57)

A radiation curing composition suitable for building a threedimensional object by a solid freeform method is disclosed. The composition includes one or more mono-functional monomers where a respective polymer has a Glass Transition Temperature higher than about 60° C., one or more di-functional oligomers where a respective polymer has a Glass Transition Temperature lower than about 40° C. and a phase separation inducing component. The phase separation may be induced during curing, resulting in a non-clear cured material having an improved impact strength and higher elongation, when compared to similar compositions without a phase separation including component.

Provisional application No. 60/188,698, filed on Mar. (60)13, 2000, provisional application No. 60/195,321, filed on Apr. 10, 2000.

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15 Claims, 3 Drawing Sheets

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#### COMPOSITIONS AND METHODS FOR USE IN THREE DIMENSIONAL MODEL PRINTING

#### CROSS REFERENCE TO RELATED APPLICATIONS

This patent application is a continuation application of U.S. patent application Ser. No. 12/963,577 filed Dec. 28, 2010 now U.S. Pat. No. 8,106,107 which will be issued as 10U.S. patent Ser. No. 12/963,577. U.S. patent application Ser. No. 12/963,577 is a continuation application of 12/342,210, filed Dec. 23, 2008, now U.S. Pat. No. 7,851,122, which in turn is a continuation application of prior U.S. patent application Ser. No. 11/518,295, filed Sep. 11, 2006, now U.S. Pat. <sup>15</sup> No. 7,479,510, which in turn is a continuation application of prior U.S. patent application Ser. No. 11/098,690, filed Apr. 5, 2005, now U.S. Pat. No. 7,183,335, which is a continuation application of prior U.S. application Ser. No. 10/424,732, filed Apr. 29, 2003, now abandoned, which is a Continuation-<sup>20</sup> in-part application of U.S. patent application Ser. No. 09/803, 108, filed Mar. 12, 2001, now U.S. Pat. No. 6,569,373, which claims priority from Provisional application No. 60/188,698, filed Mar. 13, 2000 and Provisional application No. 60/195, 321 filed Apr. 10, 2000, each of which are incorporated in its <sup>25</sup> entirety by reference herein.

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In U.S. patent application Ser. No. 09/412,618 to the Assignees of the present invention, and incorporated herein by reference, embodiments are described including an apparatus and a method for three-dimensional model printing. Some embodiments of this application describe a system and a method for printing complex three-dimensional models by using interface materials having different hardness or elasticity and mixing the interface material from each of the printing heads to control the hardness of the material forming the three-dimensional model. The construction layers of the model may be formed from interface material having a different (harder) modulus of elasticity than the material used to form the release (and support) layers, thereby allowing for the forming of complex shapes. Radiation curable inks are disclosed in, for example, U.S. Pat. Nos. 4,303,924, 5,889,084, and 5,270,368. U.S. Pat. No. 4,303,924 discloses, inter alia, radiation curable compositions for jet-drop printing containing multifunctional ethylenically unsaturated material, monofunctional ethylenically unsaturated material, a reactive synergist, a dye colorant and an oil soluble salt. U.S. Pat. No. 5,889,084 discloses, inter alia, a radiation curable ink composition for ink-jet printing which includes a cationically photoreactive epoxy or vinyl ether monomer or oligomer, a cationic photo-initiator and a coloring agent. U.S. Pat. No. 5,270,368 discloses, inter alia, a UV curable ink composition for ink-jet printing including a resin formulation having at least two acrylate components, a photo-initiator and an organic carrier. The ink compositions disclosed in these references are typically formulated for use in ink-jet printing. Compositions for ink-jet printing are typically formulated differently from compositions for building three-dimensional objects, and thus have different properties. For example, high viscosity at room temperature is a desirable property for three-dimen-<sup>35</sup> sional objects, and thus compositions for building three-dimensional objects are typically designed to have a high viscosity at room temperature. In contrast, compositions for ink-jet printing are designed to have low viscosity at room temperature in order to function well in the printing process. None of the above-mentioned references disclose compositions that are especially formulated for three-dimensional printing. Radiation curable compositions for stereolithography are disclosed in U.S. Pat. No. 5,705,316. U.S. Pat. No. 5,705,316 discloses compounds having at least one vinyl ether group, which also contain in the molecule at least one other functional group such as an epoxy or an acrylate group; compositions including these compounds; and methods of producing three-dimensional objects using these compositions. The compounds of U.S. Pat. No. 5,705,316 are complex molecules that are not readily available and thus need to be especially synthesized, which incurs additional time and costs.

#### FIELD OF THE INVENTION

The present invention relates to three-dimensional object <sup>30</sup> building in general and to methods and compositions for use in three-dimensional printing of complex structures in particular.

#### BACKGROUND OF THE INVENTION

Three-dimensional printing, which typically works by building parts in layers, is a process used for the building up of three-dimensional objects. Three-dimensional printing is relatively speedy and flexible, allowing for the production of 40 prototype parts, tooling and rapid manufacturing of threedimensional complex structures directly from a CAD file, for example.

Using three-dimensional printing may enable a manufacturer to obtain a full three-dimensional model of any proposed 45 product before tooling, thereby possibly substantially reducing the cost of tooling and leading to a better synchronization between design and manufacturing. A lower product cost and improved product quality may also be obtained.

Using three-dimensional printing also enables the direct 50 manufacturing of full three-dimensional objects, thereby substantially reducing costs and leading to a better synchronization between design, production and consumption (use). A lower product cost and improved product quality may thus also be obtained. 55

Various systems have been developed for computerized three-dimensional printing. In U.S. Pat. No. 6,259,962 to the Assignees of the present application, and incorporated herein by reference, embodiments of an apparatus and a method for three-dimensional model printing are described. The appara-60 tus according to some embodiments in this patent include a printing head having a plurality of nozzles, a dispenser connected to the printing head for selectively dispensing interface material in layers, and curing means for optionally curing each of the layers deposited. The depth of each deposited 65 layer may be controllable by selectively adjusting the output from each of the plurality of nozzles.

Thus, there is a need for simple, easily obtainable curable compositions that are specially formulated to construct a three-dimensional object. There is further a need for simple, easily obtainable curable compositions, that are specially formulated to provide support to a three-dimensional object, by forming support/and or release layers around the object during its construction. Lastly, there is a need for methods of constructing a three-dimensional object by using the above mentioned compositions.

#### SUMMARY OF THE INVENTION

The present invention relates to compositions for use in the manufacture of three-dimensional objects. The present inven-

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tion further relates to compositions for use as a support and/or release material in the manufacture of the three-dimensional objects. The present invention further relates to a method for the preparation of a three-dimensional object by three-dimensional printing, and to a three-dimensional object obtained by 5 the method.

There is thus provided, in accordance with an embodiment of the present invention, a composition for use in the manufacture of three-dimensional objects by a method of selective dispensing. The composition may include, inter alia, at least 10 one reactive component, at least one photo-initiator, at least one surface-active agent, and at least one stabilizer.

The composition has a first viscosity above 50 cps at room temperature, and a second viscosity compatible with ink-jet printers at a second temperature, wherein the second tempera 15 ture is higher than room temperature. In accordance with an embodiment of the present invention, the reactive component is an acrylic component, a molecule having one or more epoxy substituents, a molecule having one or more vinyl ether substituents, vinylcaprolac- 20 tam, vinylpyrolidone, or any combination thereof. Furthermore, in accordance with an embodiment of the present invention, the reactive component is an acrylic component. The acrylic component is an acrylic monomer, an acrylic oligomer, an acrylic crosslinker, or any combination 25 thereof. Furthermore, in accordance with an embodiment of the present invention, the reactive component may include, inter alia, an acrylic component and in addition a molecule having one or more epoxy substitutents, a molecule having one or 30 more vinyl ether substituents, vinylcaprolactam, vinylpyrolidone, or any combination thereof.

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In addition, in accordance with another embodiment of the present invention, there is thus provided a composition for use as a support and/or release material in the manufacture of three-dimensional objects by a method of selective dispensing. The composition may include, inter alia, at least one non-reactive and low toxicity compound, at least one surfaceactive agent and at least one stabilizer.

The composition has a first viscosity above 50 cps at room temperature, and a second viscosity compatible with ink-jet printers at a second temperature, wherein the second temperature is higher than room temperature.

In accordance with an embodiment of the present invention, the composition may further include, inter alia, at least one reactive component and at least one photo-initiator. The reactive component is at least one of an acrylic component, a molecule having one or more vinyl ether substituents, or the reactive component is a water miscible component that is, after curing, capable of swelling upon exposure to water or to an alkaline or acidic water solution. Furthermore, in accordance with an embodiment of the present invention the reactive component is an acrylic component. The acrylic component is an acrylic oligomer, an acrylic monomer, or a combination thereof. Furthermore, in accordance with an embodiment of the present invention, the reactive component may include, inter alia, at least one water miscible component that is, after curing, capable of swelling upon exposure to water or to an alkaline or acidic water solution. The water miscible component is preferably an acrylated urethane oligomer derivative of polyethylene glycol, a partially acrylated polyol oligomer, an acrylated oligomer having hydrophillic substituents, or any combination thereof. The hydrophilic substituents are preferably acidic substituents, amino substituents, hydroxy substituents, or any combination thereof.

Furthermore, in accordance with an embodiment of the present invention, the reactive component may include, inter alia, an acrylic component and vinylcaprolactam. Furthermore, in accordance with an embodiment of the present invention, the reactive component may include, inter alia, a molecule having one or more vinyl ether substitutents. Furthermore, in accordance with an embodiment of the present invention, the reactive component may include, inter 40 alia, a molecule having one or more epoxy substituents. Furthermore, in accordance with an embodiment of the present invention, the reactive component may include, inter alia, a molecule having one or more vinyl ether substituents, and a molecule having one or more epoxy substitutents. Furthermore, in accordance with an embodiment of the present invention, the photo-initiator is a free radical photoinitiator, a cationic photo-initiator, or any combination thereof. Furthermore, in accordance with an embodiment of the 50 present invention, the composition further includes at least one pigment and at least one dispersant. The pigment is a white pigment, an organic pigment, an inorganic pigment, a metal pigment or a combination thereof. In one embodiment, the composition further includes a dye.

Furthermore, in accordance with an embodiment of the present invention, the reactive component may include, inter alia, a molecule having one or more vinyl ether substituents. Furthermore, in accordance with an embodiment of the present invention, the non-reactive component is polyethylene glycol, methoxy polyethylene glycol, glycerol, ethoxylated polyol, or propylene glycol. Furthermore, in accordance with an embodiment of the present invention, the photo-initiator is a free radical photoinitiator, a cationic photo-initiator, or a combination thereof. Furthermore, in accordance with an embodiment of the 45 present invention, the first viscosity of the composition is greater than 80 cps. In one embodiment, the first viscosity is between 80 and 300 cps. In another embodiment, the first viscosity is around 200 cps. Furthermore, in accordance with an embodiment of the present invention, the second viscosity of the composition is lower than 20 cps at a second temperature, which is greater than 60° C. Preferably, the second viscosity is between 8 and 15 cps at the second temperature, which is greater than 60° C. 55 In one embodiment, the second viscosity is about 11 cps at a temperature around 85° C.

Furthermore, in accordance with an embodiment of the present invention, the first viscosity of the composition is greater than 80 cps. In one embodiment, the first viscosity is between 80 and 300 cps. In another embodiment, the first viscosity is around 300 cps. 60 Furthermore, in accordance with an embodiment of the present invention, the second viscosity of the composition is lower than 20 cps at a second temperature, which is greater than 60° C. Preferably, the second viscosity is between 8 and 15 cps at the second temperature, which is greater than 60° C. 65 In one embodiment, the second viscosity is about 11 cps at a temperature around 85° C.

In addition, there is thus provided, in accordance with an embodiment of the present invention, a method for preparation of a three-dimensional object by three-dimensional print-60 ing. The method according to an embodiment includes: dispensing a first interface material from a printing head, the first interface material may include, inter alia, at least one reactive component, at least one photo-initiator, at least one surface-active agent and at least one stabilizer, dispensing a 65 second interface material from the printing head, the second interface material may include, inter alia, at least one nonreactive and low toxicity compound, at least one surface-

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active agent, and at least one stabilizer, combining the first interface material and the second interface material in predetermined proportions to produce construction layers for forming the three-dimensional object.

Furthermore, in accordance with an embodiment of the 5 present invention, the reactive component of the first interface material is an acrylic component, a molecule having one or more epoxy substituents, a molecule having one or more vinyl ether substituents, vinylpyrolidone, vinylcaprolactam, or any combination thereof.

Furthermore, in accordance with an embodiment of the present invention, the reactive component of the first interface material may include, inter alia, an acrylic component. The acrylic component is an acrylic monomer, an acrylic oligomer, an acrylic crosslinker, or any combination thereof. Furthermore, in accordance with an embodiment of the present invention, the reactive component of the first interface material may include, inter alia, an acrylic component and in addition a molecule having one or more epoxy substituents, a molecule having one or more vinyl ether substituents, vinyl-20 caprolactam, vinylpyrolidone, or any combination thereof. Furthermore, in accordance with an embodiment of the present invention, the reactive component of the first interface material may include, inter alia, an acrylic component and vinylcaprolactam. Furthermore, in accordance with an embodiment of the present invention, the reactive component of the first interface material is a molecule having one or more vinyl ether substituents.

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an acrylated oligomer having hydrophillic substituents, or any combination thereof. The hydrophilic substituents are preferably acidic substituents, amino substituents, hydroxy substituents, or any combination thereof.

Furthermore, in accordance with an embodiment of the present invention, the reactive component of the second interface material may include, inter alia, a molecule having one or more vinyl ether substituents.

Furthermore, in accordance with an embodiment of the present invention, the non-reactive component is polyethylene glycol, methoxy polyethylene glycol, glycerol, ethoxylated polyol, or propylene glycol.

Furthermore, in accordance with an embodiment of the present invention, the photo-initiator of the first interface material and optionally of the second interface material is a free radical photo-initiator, a cationic photo-initiator or any combination thereof.

Furthermore, in accordance with an embodiment of the 30 present invention, the reactive component of the first interface material is a molecule having one or more epoxy substituents.

Furthermore, in accordance with an embodiment of the present invention, the reactive component of the first interface material may include, inter alia, a molecule having one or 35 more epoxy substituents, and a molecule having one or more vinyl ether substituents. Furthermore, in accordance with an embodiment of the present invention, the first interface material may further include, inter alia, at least one pigment and at least one dis- 40 persant. The pigment is a white pigment, an organic pigment, an inorganic pigment, a metal pigment or a combination thereof. In one embodiment, the first interface material may further include, inter alia, a dye.

Furthermore, in accordance with an embodiment of the present invention, the method may further include the step of irradiating or curing the second interface material.

Furthermore, in accordance with an embodiment of the present invention, the first interface material and the second interface material have a different modulus of elasticity and a 25 different strength. In one embodiment, the first interface material has a higher modulus of elasticity and a higher strength than the second interface material.

Furthermore, in accordance with an embodiment of the present invention, the method may further include the step of forming a multiplicity of support layers for supporting the object. In one embodiment, the support layers are formed by combining the first interface material and the second interface material in pre-determined proportions. In one embodiment, the support layers have the same modulus of elasticity and the same strength as the construction layers. In another embodi-

Furthermore, in accordance with an embodiment of the 45 present invention, the method may further include the step of curing the first interface material.

Furthermore, in accordance with an embodiment of the present invention, the second interface material further may include, inter alia, at least one reactive component and at least 50 one photo-initiator. The reactive component is at least one of an acrylic component, a molecule having one or more vinyl ether substituents, or the reactive component is a water miscible component that is, after curing, capable of swelling upon exposure to water or to an alkaline or acidic water 55 solution.

Furthermore, in accordance with an embodiment of the present invention the reactive component is an acrylic component. The acrylic component is an acrylic oligomer, an acrylic monomer, or a combination thereof.

ment, the support layers have a lower modulus of elasticity and a lower strength than the construction layers.

Furthermore, in accordance with an embodiment of the present invention, the method may further include the step of combining the first interface material and the second interface material in pre-determined proportions to form a multiplicity of release layers for releasing the support layers from the object. In one embodiment, the release layers have a lower modulus of elasticity and a lower strength than the construction layers and the support layers.

Furthermore, in accordance with an embodiment of the present invention, the first interface material and the second interface material each have a first viscosity at room temperature, and a second viscosity compatible with ink-jet printers at a second temperature, which may be the same or different, wherein the second temperature is higher than room temperature.

In addition, there is thus provided, in accordance with another embodiment of the present invention, a three-dimensional object comprised of a core consisting of a multiplicity of construction layers. The construction layers are prepared by combining pre-determined proportions of the first interface material and the second interface material, described hereinabove. Furthermore, in accordance with an embodiment of the 60 present invention, the object may further include a multiplicity of support layers for supporting the core. In one embodiment, the support layers are prepared by combining predetermined proportions of the first interface material and a second interface material. In one embodiment, the support layers have the same modulus of elasticity and the same strength as the construction layers. In another embodiment,

Furthermore, in accordance with an embodiment of the present invention, the reactive component may include, inter alia, at least one water miscible component that is, after curing, capable of swelling upon exposure to water or to an alkaline or acidic water solution. The water miscible compo-65 nent is preferably an acrylated urethane oligomer derivative of polyethylene glycol, a partially acrylated polyol oligomer,

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the support layers have a lower modulus of elasticity and a lower strength than the construction layers.

Furthermore, in accordance with a preferred embodiment of the present invention, the object may further include a multiplicity of release layers for releasing the support layers 5 from the core. In one embodiment, the release layers are positioned between the support layers and the construction layers. The release layers are prepared by combining predetermined proportions of the first interface material and a second interface material. In one embodiment, the release 10 layers have a lower modulus of elasticity and a lower strength than the construction layers and the support layers.

One embodiment of the present invention provides a composition suitable for building a three-dimensional object, the composition may include, inter alia, a curable component, 15 having a functional group, wherein if the functional group is a polymerizable reactive functional group, then the functional group is a (meth)acrylic functional group, a photo-initiator, a surface-active agent and a stabilizer, wherein the composition has a first viscosity of about 50-500 cps at a first temperature, 20 wherein the first temperature is ambient temperature, and a second viscosity lower than 20 cps at a second temperature wherein the second temperature is higher than the first temperature, wherein, after curing, the composition results in a solid form. One embodiment of the present invention provides a composition suitable for support in building a three-dimensional object, the composition may include, inter alia, a non-curable component, a curable component, wherein the non-curable component is not reactive with the curable component, a surface-active agent and a stabilizer, wherein the composition has a first viscosity of about 20-500 cps at a first temperature, wherein the first temperature is ambient temperature, and a second viscosity lower than 20 cps at a second temperature wherein the second temperature is higher than the first temperature, wherein, after irradiation, the composition results in a solid, a semi solid or a liquid material. One embodiment of the present invention provides a composition suitable for support in building a three-dimensional object, the composition may include, inter alia, a non-curable 40 component, a curable (meth)acrylic component, wherein the non-curable component is not reactive with the curable component, a surface-active agent, a free radical photo-initiator and a stabilizer, wherein the composition has a first viscosity of about 20-500 cps at a first temperature, wherein the first 45 temperature is ambient temperature, and a second viscosity lower than 20 cps at a second temperature wherein the second temperature is higher than the first temperature, wherein, after irradiation, the composition results in a solid, a semisolid or a liquid material. One embodiment of the present invention further provides a composition suitable for support in building a three-dimensional object, the composition may include, inter alia, at least one non-curable component, at least one curable component including a molecule having one or more epoxy substituents, 55 wherein the non-curable component is not reactive with the curable component, at least one surface-active agent, at least one cationic photo-initiator and at least one stabilizer, wherein the composition has a first viscosity of about 20-500 cps at a first temperature, wherein the first temperature is 60 ambient temperature, and a second viscosity lower than 20 cps at a second temperature wherein the second temperature is higher than the first temperature, wherein, after irradiation, the composition results in a solid, a semi solid or a liquid material.

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three-dimensional printing, the method may include the steps of dispensing a first composition suitable for building a threedimensional object from a dispenser, the first composition may include a curable component, having a functional group, wherein if the functional group is a polymerizable reactive functional group, then the functional group is a (meth)acrylic functional group, a photo-initiator, a surface-active agent, and a stabilizer, dispensing a second composition suitable for support in building a three-dimensional object from a dispenser, the second composition may include a non-curable component, a curable component, wherein the non-curable component is not reactive with the curable component, a surface-active agent and a stabilizer, combining the first composition and the second composition in pre-determined proportions to produce a multiplicity of construction layers for forming the three-dimensional object, whereby the first composition is cured resulting in a solid form, and whereby the second composition is irradiated or cured resulting in a liquid, a solid or a semi-solid form. One embodiment of the present invention further provides a three-dimensional object comprised of a multiplicity of construction layers, wherein the construction layers are prepared by combining pre-determined proportions of a first composition and a second composition according to the <sup>25</sup> invention.

#### BRIEF DESCRIPTION OF THE DRAWINGS

The present invention will be understood and appreciated more fully from the following detailed description taken in conjunction with the appended drawings in which:

FIG. 1 is a schematic illustration of an embodiment of a three-dimensional printing system;

FIG. 2 is an elevational view of a three-dimensional object, constructed in accordance with an embodiment of the present

invention; and

FIG. **3** is a schematic illustration of an embodiment of a method for the preparation of three-dimensional object by three-dimensional printing.

#### DETAILED DESCRIPTION OF THE PRESENT INVENTION

The present invention relates to compositions for use in the 45 manufacture of three-dimensional objects, and to compositions for use as support and/or release material in the manufacture of three-dimensional objects. The present invention further relates to a method for the preparation of a threedimensional object by three-dimensional printing, using the 50 above-mentioned compositions, and to a three-dimensional object obtained by the method.

The composition for use in the manufacture of the threedimensional objects may include, inter alia, at least one reactive component, at least one photo-initiator, at least one surface-active agent and at least one stabilizer. The composition may be formulated so as to be compatible for use with ink-jet printers and to have a viscosity at room temperature above 50 cps.

One embodiment of the present invention further provides a method for the preparation of a three-dimensional object by

The composition for use as a support and/or second interface material in the manufacture of the three-dimensional objects may include, inter alia, at least one non-reactive and low-toxicity component, at least one surface-active agent and at least one stabilizer. The composition may further contain at least one reactive component and at least one photo-initiator.
The composition is formulated so as to be compatible for use with ink-jet printers and to have a viscosity at room temperature above 50 cps.

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The compositions will be described in further detail below. The three-dimensional object of the present invention may be built using, for example, a three-dimensional printing system similar to embodiments of U.S. patent application Ser. No. 09/412,618, assigned to the Assignees of the present 5 application and incorporated herein by reference, although other suitable three-dimensional printers may be used. A three-dimensional printing system is shown in FIG. 1, to which reference is now made. FIG. 1 is an illustration of a three-dimensional printing system, generally designated 10, 10 which includes one or more printing heads, referenced 12, and at least two dispensers generally referenced 14 and individually referenced 14a and 14b, containing interface materials, generally referenced 16 and individually referenced 16a and 16b, respectively. Other components, and other sets of 15 printing apparatus during the printing process. components, may be used. Printing head 12 has a plurality of ink-jet type nozzles 18, through which interface materials 16a and 16b are jetted. In one embodiment of the present invention, first dispenser 14a is connected to a first set of nozzles, referenced 18a, and 20 second dispenser 14b is connected to a second set of nozzles, referenced 18b. Thus first interface material 16a is jetted through nozzles 18a, and second interface material 16b is jetted through nozzles 18b. Alternatively, in another embodiment (not shown), the three-dimensional printing system may 25 include at least two printing heads. The first printing head is connected to first dispenser 14a and is used to jet first interface material 16*a*; and the second printing head is connected to second dispenser 14b is used to jet second interface material **16***b*. The three-dimensional printing system 10 further includes a controller 20, a Computer Aided Design (CAD) system 22, curing unit 24, and optionally a positioning apparatus 1. The layers. controller 20 is coupled to the CAD system 22, curing unit 24, positioning apparatus 1, printing head 12 and each of the 35 dispensers 14. Control may be effected by other units than shown, such as one or more separate units. The three-dimensional object being produced (28) is built in layers, the depth of each layer typically being controllable by selectively adjusting the output from each of the ink-jet 40 nozzles 18. By combining or mixing materials from each of the dispensers, wherein each dispenser contains interface material having a different hardness, it is possible to adjust and control the hardness of the material forming the three-dimensional 45 object being produced. Thus, by combining the first and second interface materials being output from each of the dispensers, respectively, different parts of the three-dimensional object having a different modulus of elasticity and a different strength may be produced. As used hereinafter, the term "strength" is used as a relative term to indicate the difference in modulus of elasticity among interface materials. The strength of a material may be described, for example, by reference to its modulus of elasticity, which may be defined as: "the ratio of stress to its 55 corresponding strain under given conditions of load, for materials that deform elastically, according to Hooke's law". In accordance with one embodiment of the present invention, the first dispenser 14*a* contains a first interface material 16*a*, referred to hereinafter as the "first interface material" or 60"first composition", and the second dispenser 14b contains a second interface material 16b, referred to hereinafter as the "second interface material" or "second composition". The first interface material has a different (harder) modulus of elasticity and a greater strength than the second interface 65 material. By combining the first interface material and the second interface material, different layers of the three-dimen-

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sional object having a different modulus of elasticity and a different strength may be produced, such as, for example, a model or "construction" layer (otherwise known as a model construction), a support layer (otherwise known as a support construction) and a release layer (otherwise known as a release construction), as defined hereinbelow. In accordance with embodiments of the present invention, each layer of materials deposited by the apparatus during the printing process, may include a combination of model constructions, support constructions and/or release constructions, according to the requirements of the three-dimensional object being printed. Thus, when referring herein to construction layers, support layers and/or release layers, any or all of these may be part or parts comprising a single whole 'layer' printed by the For example, combining the first interface material and the second interface material forms a multiplicity of construction layers, which are defined as the layers constituting the threedimensional object. Multiplicity, as used hereinafter, refers to a number which is one or greater. Further, combining the first interface material and the second interface material may form a multiplicity of support layers, which are defined as the layers supporting the threedimensional object, and not constituting the three-dimensional object. Further, combining the first interface material and the second interface material may form a multiplicity of release layers, which are defined as the layers (not constituting the three-dimensional object) for separating the three-dimen-30 sional object layer from layers such as the support layers. The release layers typically have a lower modulus of elasticity and a lower strength than the construction layers and the support

In one embodiment of the present invention, the support layers are designed substantially exactly as the construction layers, and thus have substantially the same modulus of elasticity and substantially the same strength as the construction layers. In this way, the construction layers form a core, and the support layers look like the negative printing of the core. The release layers are positioned between the construction layers and the support layers, and are used to separate the construction layers from the support layers. In one embodiment of the present invention, the support layers have a lower modulus of elasticity and a lower strength than the construction layers. The support layers may be separated from the construction layers by taking advantage of their weaker properties, as will be explained in detail below. Alternatively, the support layers may be separated from the construction layers by positioning release layers between the 50 construction layers and the support layers. In order to more clearly define the present invention, reference is now made to FIG. 2, which is a three-dimensional model of a wineglass, generally referenced **30**. This threedimensional model is printed using the ink-jet type printing system of FIG. 1. combining the first interface material and the second interface material to form a multiplicity of construction layers 32 which make up wine glass 30. The construction layers 32 of wineglass 30 need to be supported externally, such as in the area referenced 34. Furthermore, an internal void, referenced **36**, needs to be formed during printing. Thus a multiplicity of support layers 38, formed by combining the first interface material and the second interface material, are printed. Furthermore, combination of the first interface material and the second interface material forms a multiplicity of release layers 40. In one embodiment of the present invention, release layers 40 are positioned between construction layers

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32 and support layers 38. Generally, release layers 40 have a different (lower) modulus of elasticity than support layers 38 and construction layers 32. Thus release layers 40 may be used to separate support layers 38 from construction layers 32.

The present invention, which will now be described in detail, provides compositions suitable for use as the first interface and as the second interface material.

The first interface material and second interface material of the present invention are especially designed and formulated 10 for building a three-dimensional object using three-dimensional printing. Accordingly, in accordance with an embodiment of the present invention, the first interface material and the second interface material each have a first viscosity at room temperature, and a second viscosity compatible with 15 ink-jet printers at a second temperature, which may be the same or different, wherein the second temperature is higher than room temperature, which is defined as about 20-30° C. In one embodiment of the present invention, the first and the second interface materials are designed to have increased 20 viscosity at room temperature, which is defined as about 20-30° C. In another embodiment, the first and second interface material have a viscosity greater than 50 cps at room temperature, In another embodiment, the viscosity may be between 80 and 300 cps. In another embodiment, the first and 25 the second interface material may have a viscosity of around 300 cps at room temperature. In one embodiment of the present invention, the first interface material and the second interface material may have a second viscosity compatible with ink-jet printing, at a second 30 temperature which may be higher than room temperature. In another embodiment, a composition compatible with ink-jet printing may have a low viscosity, for example, below 20 cps at the printing temperature, in order to function properly in the printing process. In another embodiment, the first inter- 35 face material and the second interface material, upon heating, have a viscosity preferably below 20 cps that may enable the construction of the three-dimensional object under heat. In one embodiment of the present invention, the temperature typically used to build the three-dimensional model of the 40 present invention is higher than 60° C. In another embodiment, the temperature may be about 85° C. In one embodiment of the present invention, the first and second interface materials may have a viscosity of 8-15 cps at a temperature greater than 60° C. In another embodiment, the first and 45 second interface materials may have a viscosity of 11 cps at a temperature of about 85° C. Having this viscosity, the first and second interface material in one embodiment may be distinguished from prior art formulations designed for ink-jet printing, which have low 50 viscosity at room temperature, the temperature at which the printing is normally conducted. High viscosity at room temperature is a desirable property for three-dimensional objects, a feature that is lacking in the prior art formulations. Of course, other embodiments may have other viscosities. First Interface Material

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One embodiment of the present invention provides a first interface material of the present invention may include, inter alia, at least one reactive component, at least one photoinitiator, at least one surface-active agent and at least one stabilizer.

One embodiment of the present invention provides a composition suitable for building a three-dimensional object, the composition may include, inter alia, a curable component, having a functional group, wherein if the functional group is a polymerizable reactive functional group, then the functional group is a (meth)acrylic functional group, a photo-initiator, a surface-active agent and a stabilizer, wherein the composition has a first viscosity of about 50-500 cps at a first temperature, wherein the first temperature is ambient temperature, and a second viscosity lower than 20 cps at a second temperature wherein the second temperature is higher than the first temperature, wherein, after curing, the composition results in a solid form. In one embodiment of the present invention, the first temperature is a room temperature. In another embodiment, the room temperature is between 20-30° C. In another embodiment, the first temperature is ambient temperature. In another embodiment, ambient temperature is between 10-40° C. In another embodiment, ambient temperature is between 15-35° C. In another embodiment, ambient temperature is between 20-30° C. In one embodiment of the present invention, the second temperature is higher than 40° C. In another embodiment, the second temperature is higher than 50° C. In another embodiment, the second temperature is higher than 60° C. In another embodiment, the second temperature is higher than  $70^{\circ}$  C. In one embodiment of the present invention, the curable component is a reactive component, which is able to undergo polymerization. In one embodiment of the present invention,

The first interface material (typically, the model material)

the curable component may be a (meth)acrylic monomer, a (meth)acrylic oligomer, a (meth)acrylic crosslinker, or any combination thereof.

In one embodiment of the present invention, the curable component may be a combination of a mono-functional monomer and a di-functional oligomer.

In one embodiment of the present invention, the monofunctional monomer is a high Glass Transition Temperature mono-functional monomer. In another embodiment, the difunctional oligomer is a low Glass Transition Temperature di-functional oligomer. The term Glass transition temperature (Tg) is defined as the temperature at which a polymer changes from hard and brittle to soft and pliable material.

In one embodiment of the present invention, the Glass 50 Transition Temperature of the mono-functional monomer may be higher than 60° C. In another embodiment, the Glass Transition Temperature of the mono-functional monomer may be higher than 70° C. In another embodiment, the Glass Transition Temperature of the mono-functional monomer 55 may be in the range of 70-110° C.

In one embodiment of the present invention, the Glass Transition Temperature of the di-fuctional oligomer may be lower than 40° C. In another embodiment, the Glass Transition Temperature of the di-fuctional oligomer may be lower than 30° C. In another embodiment, the Glass Transition Temperature of the di-fuctional oligomer may be in the range of 20-30° C. One embodiment of the present invention provides a composition wherein the Glass Transition Temperature of the mono-functional monomer is higher than 70° C. and wherein the Glass Transition Temperature of the di-functional oligomer is lower than 40° C.

is a composition suitable for building a three-dimensional object. The composition may be formulated to give, after curing, a solid material. In one embodiment, this invention 60 describes a composition that after curing results in a solid material, with mechanical properties that permit the building and handling of that three-dimensional object. In another embodiment, this invention provides a composition that upon curing results in a solid elastomer like material, with 65 mechanical properties that permit the building and handling of the three-dimensional object.

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In one embodiment of the present invention, the composition may include at least 20% of the high Glass Transition Temperature mono-functional monomer. In another embodiment, the composition may include at least 30% of the high Glass Transition Temperature mono-functional monomer. In 5 another embodiment, the composition may include at least 40% of the high Glass Transition Temperature mono-functional monomer. In another embodiment, the composition may include between 20-40% of the high Glass Transition Temperature mono-functional monomer. In another embodi- 10 ment, the composition may include between 30-60% of the high Glass Transition Temperature mono-functional monomer. In one embodiment of the present invention, the composition may include about 20% of the low Glass Transition 15 Temperature di-functional oligomers. In another embodiment, the composition may include about 40% of the low Glass Transition Temperature di-functional oligomers. In another embodiment, the composition may include between 20-40% of the low Glass Transition Temperature di-func- 20 tional oligomers. In another embodiment, the composition may include at least 20% of the low Glass Transition Temperature di-functional oligomer. In another embodiment, the composition may include not more than 40% of the low Glass Transition Temperature di-functional oligomer. In one embodiment of the present invention, the composition may include at least 40% of the high Glass Transition Temperature mono-functional monomers and at least 20% of the low Glass Transition Temperature di-functional oligomer. In one embodiment of the present invention, the composi- 30 tion may include at least 20% of the high Glass Transition Temperature mono-functional monomers and not more than 40% of the low Glass Transition Temperature di-functional oligomer.

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more vinyl ether substituents. In one embodiment of the present invention, the concentration of component that includes a molecule having one or more vinyl ether substituents is in the range of 10-30%. In another embodiment, the concentration is 15-20%. In another embodiment, the concentration is 15%. Of course, other concentrations, and other ranges, can be used. Conventional vinyl ether monomers and oligomers which have at least vinyl ether group are suitable. Examples of vinyl ethers are ethyl vinyl ether, propyl vinyl ether, isobutyl vinyl ether, cyclohexyl vinyl ether, 2-ethylhexyl vinyl ether, butyl vinyl ether, ethyleneglocol monovinyl ether, diethyleneglycol divinyl ether, butane diol divinyl ether, hexane diol divinyl ether, cyclohexane dimethanol monovinyl ether and the like. An example of a vinyl ether for the present invention is 1,4 cyclohexane dimethanol divinyl ether, marketed by ISP under the trade name CHVE. In one embodiment of the present invention, the first interface material may also include a curable component which is a molecule having one or more epoxy substituents. In one embodiment of the present invention, conventional epoxy monomers and oligomers which have at least one oxirane moiety may be used. Non-limiting examples of suitable epoxy containing molecules are displayed in Table 1 below <sup>25</sup> (note other suppliers may be used for suitable materials):

An acrylic monomer is a functional acrylated molecule 35

#### TABLE 1

| Examples of epoxy-containing curable component |                                                                    |                      |  |  |  |  |
|------------------------------------------------|--------------------------------------------------------------------|----------------------|--|--|--|--|
| Trade Name                                     | Type of Material                                                   | Supplier             |  |  |  |  |
| ERL-4299 or<br>UVR-6128                        | Bis-(3,4 cyclohexylmethyl) adipate                                 | Union Carbide        |  |  |  |  |
| UVR-6105 and<br>UVR-6110                       | 3,4-epoxy cyclohexylmethyl-3,4-<br>epoxycyclohexyl carboxylate     | Union Carbide        |  |  |  |  |
| D.E.R 732                                      | Aliphatic epoxy, Polyglycol<br>diglycidyl ether                    | Dow chemicals        |  |  |  |  |
| Vinylcyclohexene<br>Monoxide                   | 1,2 epoxy-4-vinylcyclohexane                                       | Union Carbide        |  |  |  |  |
| D.E.N. 431                                     | Epoxy novolac resin                                                | Dow corning          |  |  |  |  |
| UVR-6216                                       | 1,2-epoxy hexadecane                                               | Union Carbide        |  |  |  |  |
| UVI-6100                                       | Cycloaliphatic epoxide diluent                                     | Union Carbide        |  |  |  |  |
| Vikoflex 7170                                  | Fullyl epoxidized soy bean oil                                     | Elf Atochem,<br>INC. |  |  |  |  |
| ERL-4221D                                      | 3,4-epoxy cyclohexylmethyl<br>3,4-epoxy cyclohexane<br>carboxylate | Union Carbide        |  |  |  |  |

which may be, for example, esters of acrylic acid and methacrylic acid. Momoners may be mono-functional or multifunctional (for example, di-, tri-, tetra-functional, and others). An example of an acrylic mono-functional monomer for the present invention is phenoxyethyl acrylate, marketed by Sar- 40 tomer under the trade name SR-339. An example of an acrylic di-functional monomer is propoxylated (2) neopentyl glycol diacrylate, marketed by Sartomer under the trade name SR-9003.

An acrylic oligomer is a functional acrylated molecule 45 which may be, for example, polyesters of acrylic acid and methacrylic acid. Other examples of acrylic oligomers are the classes of urethane acrylates and urethane methacrylates. Urethane-acrylates are manufactured from aliphatic or aromatic or cycloaliphatic diisocyanates or polyisocyanates and 50 hydroxyl-containing acrylic acid esters. An example is a urethane-acrylate oligomer marketed by Cognis under the trade name Photomer-6010.

An acrylic crosslinker is a molecule which may provide enhanced crosslinking density. Examples of such resins are 55 Ditrimethylolpropane Tetra-acrylate (DiTMPTTA), Pentaerythitol Tetra-acrylate (TETTA), Dipentaerythitol Pentaacrylate (DiPEP). In one embodiment of the present invention, the composition may further includes, inter alia, a curable component, which is a molecule having one or more 60 epoxy substituents, a molecule having one or more vinyl ether substituents, vinylcaprolactam, vinylpyrolidone, or any combination thereof. In one embodiment of the present invention, the composition may further include, inter alia, vinylcaprolactam. Other curable components may also be used. The first interface material may also include a curable component which is, for example, a molecule having one or

In one embodiment of the present invention, the first interface material may include any combination of an acrylic component as defined hereinabove, a molecule having one or more epoxy substituents as defined hereinabove, a molecule having one or more vinyl ether substituents as defined hereinabove, vinylcaprolactam and vinylpyrolidone.

In one embodiment of the present invention, the curable component of the first interface material includes, inter alia, an acrylic monomer, an acrylic oligomer, an acrylic crosslinker and vinylcaprolactam. In another embodiment, the curable component includes an acrylic component as defined hereinabove and a molecule having one or more epoxy substituents as defined hereinabove. In another embodiment, the curable component of the first interface material includes an acrylic component as defined hereinabove and a molecule having one or more vinyl ether substituents as defined hereinabove. In another embodiment, the curable component in the first interface material includes a 65 molecule having one or more vinyl ether substituents as defined hereinabove, and a molecule having one or more epoxy substituents as defined hereinabove.

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The photo-initiator of the first interface material and of the second interface material may be the same or different, and is a free radical photo-initiator, a cationic photo-initiator, or any combination thereof.

The free radical photo-initiator may be any compound that 5 produces a free radical on exposure to radiation such as ultraviolet or visible radiation and thereby initiates a polymerization reaction. Non-limiting xamples of some suitable photoinitiators include benzophenones (aromatic ketones) such as benzophenone, methyl benzophenone, Michler's ketone and 10 xanthones; acylphosphine oxide type photo-initiators such as 2,4,6-trimethylbenzolydiphenyl phosphine oxide (TMPO), 2,4,6-trimethylbenzoylethoxyphenyl phosphine oxide (TEPO), and bisacylphosphine oxides (BAPO's); benzoins and bezoin alkyl ethers such as benzoin, benzoin methyl ether 15 and benzoin isopropyl ether and the like. Examples of photoinitiators are alpha-amino ketone, marketed by Ciba Specialties Chemicals Inc. (Ciba) under the trade name Irgacure 907, and bisacylphosphine oxide (BAPO's), marketed by Ciba under the trade name 1-819. The free-radical photo-initiator may be used alone or in combination with a co-initiator. Co-initiators are used with initiators that need a second molecule to produce a radical that is active in the UV-systems. Benzophenone is an example of a photoinitiator that requires a second molecule, such as an 25 amine, to produce a curable radical. After absorbing radiation, benzophenone reacts with a ternary amine by hydrogen abstraction, to generate an alpha-amino radical which initiates polymerization of acrylates. Non-limiting example of a class of co-initiators are alkanolamines such as triethylamine, 30 methyldiethanolamine and triethanolamine. Suitable cationic photo-initiators for the present invention include compounds which form aprotic acids or Bronstead acids upon exposure to ultraviolet and/or visible light sufficient to initiate polymerization. The photo-initiator used may 35 be a single compound, a mixture of two or more active compounds, or a combination of two or more different compounds, i.e. co-initiators. Non-limiting examples of suitable cationic photo-initiators are aryldiazonium salts, diaryliodonium salts, triarylsulphonium salts, triarylselenonium salts 40 and the like. In one embodiment, a cationic photo-initiator for the present invention may be a mixture of triarylsolfonium hexafluoroantimonate salts marketed by Union Carbide as UVI-6974. In one embodiment of the present invention, the composi- 45 tion suitable for building a three-dimensional object may further include a curable compound, which is a sulfur-containing component. In one embodiment of the present invention, the sulfur containing component is beta mercaptopropionate, mercaptoacetate, alkane thiols or any combination 50 thereof. The addition of sulfur-containing components may significantly enhance the composition reactivity. At levels of about 5% of sulfur-containing component a significant reactivity enhancement is achieved. The mechanical properties of the composition may be determined depending on the sulfurcontaining component used. The reactivity enhancement achieved by the use of sulfur-containing component, enables the incorporation in the polymerization reaction of non sulfur-containing components, which would not easily polymerize otherwise. Molecules having unsaturated double bonds, 60 for example, low molecular weight polybuthadiene, are polymerized in the claimed compositions when contained an appropriate sulfur-containing component. For example, a basic composition will contain 15% low molecular weight unsaturated molecule, 5% sulfur-containing component, 15% 65 mono-functional monomer, 15% di-functional monomer and the rest other curable components according to the intended

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photopolymer properties. An example of a sulfur-containing component for the present invention is trimethylolpropane tri(3-mercaptopropionate), manufactured by BRUNO BOCK Chemische Fabrik GMBH & CO. Other suitable substances may be used.

In one embodiment of the present invention, the composition suitable for building a three-dimensional object, further includes, inter alia, a low molecular weight polymer. An example of a low molecular weight polymer for the present invention is Styrene-Butadiene-Methacrylate block copolymers (KRATON D), manufactured by Dow Corning. Other suitable substances may be used.

In one embodiment of the present invention, the composition suitable for building a three-dimensional object, further includes, inter alia, a filler.

The term filler is defined as an inert material added to a polymer, a polymer composition or other material to modify their properties and/or to adjust quality of the end products. The filler may be an inorganic particle, for example calcium carbonate, silica and clay. Of course other filler substances may be used.

Fillers may be introduced in to polymer compositions in order to reduce shrinkage during polymerization or during cooling, for example to reduce the coefficient of thermal expansion, increase strength, increase thermal stability reduce cost and/or adopt rheological properties. The use of standard fillers has also some drawbacks such as reduction of elasticity and an increase in viscosity. Additionally, large diameter fillers (>5 micron) are not appropriate for ink-jet applications.

Nano-particles fillers are especially useful in applications requiring low viscosity such as ink-jet applications. Compositions containing as much as 30% nano-particle fillers are feasible, whereas the same concentration of more standard and higher diameter fillers (~>1 micron) produce at such concentration viscosities which are too high for ink-jet applications. In one embodiment of the present invention, the nano-particle filler containing composition is clear. The composition is clear (e.g. transparent) since it contains no visual fillers. In contrast, compositions containing more standard and higher diameter visible fillers (~>1 micron), are not clear. In one embodiment of the present invention, the composition optionally may contain pigments. In another embodiment, the pigment concentration may be lower than 35%. In another embodiment, the pigment concentration may be lower than 15%. In one embodiment of the present invention, the filler may include particles such as particles having an average diameter of less than 100 nm. In another embodiment, the filler may include particles having a diameter in the range of 10-100 nm. In another embodiment, the filler may include particles having a diameter in the range of 20-80 nm. In another embodiment, the filler may include particles having a diameter in the range of 10-50 nm. In another embodiment, the filler may include particles having a diameter smaller than 10 nm. Examples of fillers that may be used in the composition are HIGHLINK OG (particle size spanning between 9 nm to 50 nm), manufactured by Clariant, and NANOCRYL (particle size below 50 nm), manufactured by Hanse Chemie. Other suitable substances may be used. It was discovered that phase separation may be induced during the radiation curing process of the present method. In one embodiment of the present invention, the phase separation may produce a clear material, which may have improved impact-resistance. This composition, upon bending develops micro-cracks, before breaking. These micro-cracks can easily be distinguished due to the whitening of the stress area or

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stress line. In another embodiment, the phase separation results in a non-clear cured material. It was discovered that certain combinations of UV curable components induce phase separation during curing. Such compositions are clear before curing and may be clear, hazy or opaque after curing. Such compositions have an improved impact strength and higher elongation, when compared to similar compositions, which do not show such phase separation. For example, it was discovered that the addition of some silicon containing oligomers, at levels as low as 5%, to the above described com- 10 position, may already create a substance which induces such phase separation. An example of a silicon acrylated molecule is Ebecryl 350, manufactured by UCB Chemicals. Of course other substances may be used. position further includes a phase separation inducing component. In another embodiment, the phase separation inducing component is a silicon oligomer. In another embodiment, the concentration of the silicon oligomer is at least 5%. In one embodiment of the present invention, phase separa-20 tion may be induced during curing, resulting in a non-clear cured material. Certain combinations of UV curable composition suffer a phase separation process during curing. Such compositions are clear before curing and hazy to white after curing. Such compositions have an improved impact strength 25 and higher elongation, when compared to similar compositions, which do not suffer from such phase separation. For example, the addition of some silicon containing oligomers, at levels as low as 5%, to the above described composition, may create a substance which suffers from such phase sepa- 30 ration.

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face material to permit the use of the formulation at high temperature, for example around 85° C., without causing thermal polymerization.

In one embodiment of the present invention, the composition may further include, inter alia, at least one pigment and at least one dispersant. In one embodiment of the present invention, the pigment may be a white pigment. In another embodiment, the pigment may be an organic pigment. In another embodiment, the pigment may be an inorganic pigment. In another embodiment, the pigment may be a metal pigment or a combination thereof. In one embodiment of the present invention, the composition may further include, inter alia, a dye. An example of a white pigment for the present invention is organic treated titanium dioxide, marketed by Kemira Pig-One embodiment of the present invention provides a com- 15 ments under the trade name UV TITAN M160 VEG. An example of an organic pigment for the present invention is an organic pigment marketed by Elementis Specialities under the trade name Tint Aid PC 9703. Examples of dispersants for the present invention are dispersants including a copolymer with acidic groups marketed by Byk Chemie under the trade name Disperbyk 110, and a dispersant including a high molecular weight block copolymer with pigment affinic groups, marketed by Byk Chemie under the trade name Disperbyk 163. Furthermore, in one embodiment of the present invention, combinations of white pigments and dyes are used to prepare colored resins. In such combinations, the white pigment may have at least a double task: 1) to impart opacity; and 2) to shield the dye from UV radiation, to prevent bleaching of the resin. Thus, in accordance with one embodiment of the present invention, the first interface material further includes a dye. The dye may be chosen so as not to interfere with the curing efficiency of the formulation of the first interface material. The dye may be any of a broad class of solvent soluble dyes. Some non-limiting examples are azo dyes 35 which are yellow, orange, brown and red; anthraquinone and

In one embodiment of the present invention, the first viscosity is about 80-500 cps. In another embodiment, the first viscosity is about 300 cps. Of course, compositions having other viscosities may be used. In one embodiment of the present invention, the second viscosity is lower than 20 cps and wherein the second temperature is higher than 60° C. In another embodiment, the second viscosity is between 10 and 17 cps and wherein the second temperature is higher than 60° C. In another embodi- 40 ment, the second viscosity is between 10 and 17 cps and wherein the second temperature is about 70-110° C. In another embodiment, the second viscosity is between 12 and 15 cps and wherein the second temperature is about 70-90° C. Of course, compositions having other viscosities may be 45 used. Other components of the first interface material and the second interface material of the present invention are surfaceactive agents and inhibitors (typically, thermal stabilizers). A surface-active agent may be used to reduce the surface tension 50 of the formulation to the value required for jetting or for printing process, which is typically around 30 dyne/cm. An example of a surface-active agent for the present invention is silicone surface additive, marketed by Byk Chemie under the trade name Byk 307 Inhibitors may be employed in the for- 55 mulations of the first interface material and the second inter-

triarylmethane dyes which are green and blue; and azine dye which is black. An example of a dye for the present invention is Solvent Red 127, marketed by Spectra Colors Corp. under the trade name Spectrasol RED BLG.

The relative proportions of the different components of the first interface material may vary. In one embodiment of the present invention, the first interface material includes the following components: 50% acrylic oligomer(s), 30% acrylic monomer(s), 15% acrylic crosslinker, 2% photoinitiator, surface active agent, pigments, and stabilizers. Of course, other compositions may be used.

Non-limiting examples of formulations of the first interface material are provided hereinbelow in Tables 2-4, to which reference is now made. Tables 2 and 3 illustrate examples of possible formulations of the first interface material. Table 4 illustrates examples of colored formulations, which include pigments, dispersants and dyes, as defined hereinabove. To any of the examples in Tables 2 and 3 may be added the combination of the colorants of Table 4. The individual substances, suppliers, combinations, etc., are given by way of example only.

TABLE 2

Examples of Characteristic Formulation Components of First Interface Material

| #           | Trade Name                        | Chemical Type                                                                              | Function in the formulation         | Supplier                       |
|-------------|-----------------------------------|--------------------------------------------------------------------------------------------|-------------------------------------|--------------------------------|
| A<br>B<br>C | Photomer-6010<br>SR-339<br>SR-351 | Urethane Acrylate Oligomer<br>Phenoxy ethyl Acrylate<br>Trimethylol<br>propane triacrylate | Oligomer<br>monomer<br>Cross-linker | Cognis<br>Sartomer<br>Sartomer |

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TABLE 2-continued

#### Examples of Characteristic Formulation Components of First Interface Material

| # | Trade Name                                               | Chemical Type                                                           | Function in the formulation     | Supplier                                           |
|---|----------------------------------------------------------|-------------------------------------------------------------------------|---------------------------------|----------------------------------------------------|
| D | Irgacure 907                                             | alpha-Amino Ketone                                                      | Free radical<br>photo-initiator | Ciba<br>Specialties<br>Chemical Inc.               |
| Е | BP                                                       | Benzophenone                                                            | Free radical photo-initiator    | Satomer                                            |
| F | Triethanol<br>Amine                                      | 1. Ternary Amine                                                        | Free radical<br>Coinitiator     | Sigma                                              |
| G | Byk 307                                                  | Silicone Surface Additive                                               | Surface agent                   | Byk Chemie                                         |
| Η | MEHQ                                                     | 4-Methoxy phenol                                                        | Inhibitor                       | Sigma                                              |
| Ι | Cyracure<br>UVR-6110                                     | 3,4 Epoxycyclohexylmethyl-<br>3,4-epoxycyclohexylcarboxylate            | Epoxy<br>oligomer               | Union Carbide                                      |
| J | UVI-6974                                                 | Mixed Triarylsulfonium<br>Hexafluoroantimonate Salts                    | Cationic<br>photo-initiator     | Union Carbide                                      |
| Κ | CHVE                                                     | 1,4-cyclohexane dimethanol divinyl ether                                | Vinyl Ether<br>Monomer          | ISP                                                |
| L | UV TITAN<br>M160 VEG                                     | Organic Treated Titanium<br>Dioxide                                     | White pigment                   | KEMIRA<br>PIGMENTS                                 |
| Μ | Disperbyk 110                                            | Copolimer with acidic groups                                            | Pigment<br>Dispersant           | Byk Chemie                                         |
| Ν | Spectrasol<br>RED BLG                                    | Solvent Red 127                                                         | Dye                             | Spectra<br>Colors Corp.                            |
| 0 | Tint Aid<br>PC 9703                                      | Organic pigment                                                         | Organic<br>pigment              | Elementis<br>Specialties                           |
| Ρ | Disperbyk 163                                            | High molecular weight block<br>copolymer with pigment affinic<br>groups | Pigment<br>Dispersant           | Byk Chemie                                         |
| Q | V-Cap                                                    | Vinylcaprolactam                                                        | Monomer                         | ISP                                                |
| R | V-Pyrol                                                  | Vinylpyrolidone                                                         | Monomer                         | ISP                                                |
| S | Silicon<br>acrylated<br>oligomer                         | Ebecryl 350                                                             | Phase<br>separation<br>promoter | UCB<br>Chemicals                                   |
| Τ | Trimethylol<br>propane<br>tri(3-mercapto-<br>propionate) | Sulfur-containing compound                                              | Crosslinker                     | BRUNO<br>BOCK<br>Chemische<br>Fabrik<br>HMBH & CO. |

TABLE 3

| Examples of Possible Formulation Compositions of First Interface Material |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
|---------------------------------------------------------------------------|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|
| Example                                                                   | А | В | С | D | Е | F | G | Η | Ι | J | Κ | Q | R | S | Т |
| 1                                                                         | Х | Х | Х | Х |   |   | Х | Х |   |   |   |   |   |   |   |
| 2                                                                         | Х | Х |   | Х |   |   | Х | Х |   |   |   |   |   |   |   |
| 3                                                                         | Х |   | Х | Х |   |   | Х | Х |   |   |   |   |   |   |   |
| 4                                                                         |   | Х | Х | Х |   |   | Х | Х |   |   |   |   |   |   |   |
| 5                                                                         | Х | Х | Х |   | Х | Х | Х | Х |   |   |   |   |   |   |   |
| 6                                                                         | Х | Х |   |   | Х | Х | Х | Х |   |   |   |   |   |   |   |
| 7                                                                         | Х |   | Х |   | Х | Х | Х | Х |   |   |   |   |   |   |   |
| 8                                                                         |   | Х | Х |   | Х | Х | Х | Х |   |   |   |   |   |   |   |
| 9                                                                         | Х |   | Х | Х |   |   | Х | Х |   |   | Х |   |   |   |   |
| 10                                                                        | Х |   | Х |   | Х | Х | Х | Х |   |   | Х |   |   |   |   |
| 11                                                                        |   |   |   |   |   |   | Х | Х | Х | Х | Х |   |   |   |   |
| 12                                                                        |   | Х | Х | Х |   |   | Х | Х | Х | Х |   |   |   |   |   |
| 13                                                                        | Х | Х | Х | Х | Х | Х | Х | Х | Х | Х | Х |   |   |   |   |
| 14                                                                        | Х | Х | Х | Х |   |   | Х | Х |   |   |   | Х |   |   |   |
| 15                                                                        | Х | Х | Х | Х |   |   | Х | Х |   |   |   |   | Х |   |   |
| 16                                                                        | Х | Х | Х | Х |   |   | Х |   |   |   |   | Х |   | Х |   |
| 17                                                                        | Х | Х | Х | Х |   |   | Х |   |   |   |   | Х |   |   | Х |

|       |                                                              |   |   | TABI | LE 4-con | tinued               |             |              |            |               |            |       |
|-------|--------------------------------------------------------------|---|---|------|----------|----------------------|-------------|--------------|------------|---------------|------------|-------|
| Exam  | Examples of colored formulations of first interface material |   |   |      |          | <b>—</b> 60 <b>—</b> | Examples of | f colored fo | rmulations | of first inte | rface mate | erial |
| Examp | ble L                                                        | М | Ν | Ο    | Р        |                      | Example     | L            | М          | Ν             | Ο          | Р     |
| 16    | Х                                                            | Х |   |      |          |                      | 19          |              |            |               | Х          | Х     |
| 17    | Х                                                            | Х | Х |      |          | 65                   | 20          |              |            | Х             | Х          | Х     |
| 18    | Х                                                            | Х |   | Х    | Х        |                      |             |              |            |               |            |       |

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In one embodiment of the present invention, the formulation of the first interface material is presented in entry No. 14 of Table No. 3. According one embodiment of the present invention, the first interface material includes:

an acrylic oligomer, which may be any acrylic oligomer as 5 defined hereinabove, and which may be an urethane acrylate oligomer;

an acrylic monomer, which may be any acrylic monomer as defined hereinabove, and which may be phenoxy ethyl acrylate;

an acrylic crosslinker, which may be any acrylic crosslinker as defined hereinabove, and which may be trimethylol propane triacrylate;

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in a liquid material. In another embodiment, the composition results in a solid material that is formulated as a weak curable material. In another embodiment, upon irradiation, the composition results in a material that is capable of swelling in water or in alkaline or acidic water. Thus, when irradiated, the second interface material swells and almost breaks upon exposure to water, with minimum manual work required.

In one embodiment of the present invention, the second interface material is formulated so as to permit fast, easy and 10 efficient removal of the second interface material and cleaning of the three-dimensional model from its support.

In one embodiment of the second invention, the curable component is a reactive component. In another embodiment of the present invention, the reactive component can undergo polymerization. According to one embodiment, the second interface material is formulated as a curable composition that is capable of solidifying upon curing. In one embodiment of the second invention, the curable components may be similar to those used in the first interface material, but chosen spe-20 cifically to give a hydrophillic cured resin, with weak mechanical properties. Thus, upon curing, a solid composition is formed that is weak and can be easily pulverized for example by hand or using water. In one embodiment of the present invention, the curable component may be a (meth)acrylic component. In another embodiment, the (meth)acrylic component may be a (meth) acrylic monomer. In another embodiment, the (meth)acrylic component may be a (meth)acrylic oligomer. In another embodiment, the (meth)acrylic component may be a (meth) acrylic crosslinker. In another embodiment, the (meth) acrylic component may be any combination of a (meth) acrylic monomer, a (meth)acrylic oligomer and a (meth) acrylic crsosslinker. In one embodiment of the present invention, the composidetergent solution and 2) a solid or semi-solid material that is 35 tion may further include, inter alia, at least one photo-initiator. In one embodiment of the present invention, the photoinitiator may a free radical photo-initiator, a cationic photoinitiator, or any combination thereof. The photo-initiator may be any photo-initiator, as defined above. One embodiment of the present invention provides a composition suitable for support in building a three-dimensional object, the composition may include, inter alia, a non-curable component, a curable (meth)acrylic component, wherein the non-curable component is not reactive with the curable com-45 ponent, a surface-active agent, a free radical photo-initiator and a stabilizer, wherein the composition has a first viscosity of about 20-500 cps at a first temperature, wherein the first temperature is ambient temperature, and a second viscosity lower than 20 cps at a second temperature wherein the second temperature is higher than the first temperature, wherein, after irradiation, the composition results in a solid, a semisolid or a liquid material. In one embodiment of the present invention, the composition may further include, inter alia, water. In one embodiment of the present invention, the composition further includes a water miscible component that is, after irradiation or curing, capable of dissolving or swelling upon exposure to water, to an alkaline or acidic water solution or to water detergent solution. In another embodiment, the water miscible component is a (meth)acrylated urethane oligomer derivative of polyethylene glycol, a partially (meth)acrylated polyol oligomer, a (meth)acrylated oligomer having hydrophillic substituents, polyethylene glycol mono or di (meth)acrylated, acrylamide, Acryloylmorpholine (ACMO) or any combination thereof. In another embodiment, the hydrophilic substituents are acidic substituents, amino substituents, hydroxy substituents, ionic substituents or any combination thereof.

a radical photo-initiator, which may be any radical photoinitiator as defined hereinabove, and which may be alpha-15 amino ketone;

a surface agent, which may be a silicone surface additive; an inhibitor, which may be 4-methoxyphenol; and vinylcaprolactam.

Second Interface Material

The second interface material (in one embodiment, the support material) is a composition typically formulated to support the building of a three-dimensional object. In one embodiment of the present invention, the second interface material is formulated to form a release layer to permit a 25 manual easy separation or cleaning of the three-dimensional object from its support.

In one embodiment of the present invention, the second interface material may be one of two different principle kinds: 1) a liquid material lacking any curable groups that remains 30 liquid even after irradiation. In one embodiment, the liquid is water miscible and is easily washed out by water, or with other material. In another embodiment the liquid is non water-miscible and is easily washed out by water or by a water formulated as a weak curable material. The solid or semisolid material, when cured, may be capable of swelling in water or in alkaline or acidic water or water detergent solution. Thus, when cured, the second interface material may swell and almost break upon exposure to water, or in alkaline 40 or acidic water or water detergent solution, with minimum manual work required. In both cases the second interface material is formulated so as to permit fast, easy and efficient removal of the second interface material and cleaning of the three-dimensional model from its support.

In one embodiment, the second interface material of the present invention may include, inter alia, at least one nonreactive and low toxicity compound, at least one surfaceactive agent and at least one stabilizer.

One embodiment of the present invention provides a com- 50 position suitable for support in building a three-dimensional object, the composition may include, inter alia, a non-curable component, a curable component, wherein the non-curable component is not reactive with the curable component, a surface-active agent, and a stabilizer, wherein the composi- 55 tion has a first viscosity of about 20-500 cps at a first temperature, wherein the first temperature is ambient temperature, and a second viscosity lower than 20 cps at a second temperature wherein the second temperature is higher than the first temperature, wherein, after irradiation, the composi- 60 tion results in a semi solid material. Of course, compositions having other viscosities may be used. In one embodiment of the present invention, the composition suitable for support in building a three-dimensional object, after irradiation, may result in a semi-solid material. In 65 another embodiment, the semi-solid material may be gel type material. In another embodiment, the composition may result

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Non-limiting examples of acrylic components for use in the second interface material of the present invention are polyethylene glycol monoacrylate, marketed by Laporte under the trade name Bisomer PEA6, polyethylene glycol diacrylate, marketed by Sartomer under the trade name 5 SR-610, methoxypolyethyleneglycole 550 monomethacrylate, and the like.

In one embodiment of the present invention, the curable component of the second interface material may be a water miscible component that is, after curing, capable of swelling 10 upon exposure to water or to an alkaline or acidic water solution. Non-limiting examples of water miscible components for the present invention are an acrylated urethane oligomer derivative of polyethylene glycol-polyethylene glycol ure than e diacrylate, a partially acrylated polyol oligomer, an 15 acrylated oligomer having hydrophillic substituents, or any combination thereof. The hydrophilic substituents are acidic substituents, amino substituents, hydroxy substituents, or any combination thereof. An example of an acrylated monomer with hydrophillic substituents is betha-carboxyethyl acrylate, 20 which contains acidic substituents. In one embodiment of the present invention, the curable component of the second interface material may also be a molecule having one or more vinyl ether substituents, which may be any of the compounds as defined hereinabove. In one 25 embodiment of the present invention, the concentration of component that includes a molecule having one or more vinyl ether substituents is in the range of 10-30%. In another embodiment, the concentration is 15-20%. In another embodiment, the concentration is 15%. Other concentrations may also be used. An example of vinyl ether for the second interface material is 1,4-cyclohexane dimethanol divinyl ether, marketed by ISP under the trade name CHVE. Other molecules having one or more vinyl ether substituents may be used. In one embodiment of the present invention, the curable component of the second interface material is an acrylic oligomer. In another embodiment, the curable component of the second interface material is a combination of an acrylic component as defined hereinabove and a water miscible compo- 40 rial. nent as defined hereinabove. In another embodiment, the curable component of the present invention is a combination of an acrylic component as defined hereinabove and a molecule having one or more vinyl ether substituents, as defined hereinabove. In another embodiment, the curable component 45 of the present invention is a combination of a water miscible component as defined hereinabove, and a molecule having one or more vinyl ether substituents, as defined hereinabove. Other combinations may also be used. In one embodiment of the present invention, the composi- 50 tion further includes, inter alia, a sulfur-containing component. In another embodiment, the sulfur containing component is beta mercaptopropionate, mercaptoacetate, alkane thiols or any combination thereof. The sulfur-containing component may be any sulfur-containing component, as 55 defined above.

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rial. High water diffusion rate is desirable in order to minimize the time needed for the water cleaning process of the three-dimensional model. Non-limiting examples of non-curable components for the present invention are polyethylene glycol marketed by Aldrich under the trade name PEG 400, methoxypolyethylene glycol marketed by Aldrich under the trade name methoxycarbowax 500 and 1000, propylene glycol and paraffin oil. Other examples are ethoxylated polyols and glycerol.

In one embodiment of the present invention, the second interface material is formulated as a liquid. The liquid formulation is a non-curable composition that remains liquid even after radiation exposure. Thus, the liquid formulation includes non-reactive components and does not include reactive components that are capable upon solidifying upon curing. In one embodiment of the present invention, the material may be water miscible, and may easily be washed out with water. In one embodiment of the present invention, the non-curable component is polyethylene glycol, methoxypolyethylene glycol, glycerol, ethoxylated polyol, propylene glycol or any combination thereof. In another embodiment, the noncurable component is a non-water miscible compound. In another embodiment, the non-water miscible compound is paraffin oil. Other non-curable substances may be used. One embodiment of the present invention further provides a composition suitable for support in building a three-dimensional object, the composition may include, inter alia, at least one non-curable component, at least one curable component including, inter alia, a molecule having one or more epoxy substituents, wherein the non-curable component is not reactive with the curable component, at least one surface-active agent, at least one cationic photo-initiator and at least one stabilizer, wherein the composition has a first viscosity of 35 about 20-500 cps at a first temperature, wherein the first temperature is ambient temperature, and a second viscosity lower than 20 cps at a second temperature wherein the second temperature is higher than the first temperature, wherein, after irradiation, the composition results in a semi solid mate-In one embodiment of the present invention, the first temperature is a room temperature. In another embodiment, the room temperature is between 20-30° C. In another embodiment, the first temperature is ambient temperature. In another embodiment, ambient temperature is between 10-40° C. In another embodiment, ambient temperature is between 15-35° C. In another embodiment, ambient temperature is between 20-30° C. In one embodiment of the present invention, the second temperature is higher than 40° C. In another embodiment, the second temperature is higher than 50° C. In another embodiment, the second temperature is higher than 60° C. In another embodiment, the second temperature is higher than  $70^{\circ}$  C. Besides swelling, another characteristic of the support upon exposure to water or to an alkaline or acidic water or water detergent solution may be the ability to break down during exposure to water or to an alkaline or acidic water solution. In one embodiment of the present invention, because the second interface material is made of hydrophillic components, during the swelling process, internal forces appear and cause fractures and breakdown of the cured second interface material. In addition, the second interface material may be at least partially water-soluble. At least part of the second interface material is may be completely water soluble/miscible. During the removal of the support and/or release layers, the water soluble/miscible components are extracted out with water.

In one embodiment of the present invention, the non-cur-

able component of the second interface material is a noncurable component. In another embodiment the non-curable component is non-polymerizing component. In another 60 embodiment, the non-curable component is a low toxicity compound. In another embodiment, the non-curable component is a water miscible one. In another embodiment, the non-curable component is a non-water miscible one. In one embodiment of the present invention, the non-curable com-65 ponent is chosen to enhance the water-swelling rate, and to reduce the mechanical strength of the second interface mate-

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In addition, in one embodiment of the present invention, the second interface material liberates bubbles upon exposure to water or to an alkaline water or acidic water solution. The bubbles are intended to help in the process of removal of the support and/or release layers from the construction layers.

In one embodiment of the present invention, the bubbles may be liberated by a bubble releasing substance (BRS) that is present in the water solution that is used to clean out the three-dimensional object. Such a substance may be a carbonate or bicarbonate, for example sodium bicarbonate (SBC). <sup>10</sup> During the swelling process, at least part of the SBC is introduced or absorbed into the second interface material, where it is transformed into carbon dioxide gas (CO<sub>2</sub>) and a watersoluble salt. The trigger for the production of  $CO_2$  may be the reaction of the SBC with an acid functionality present in the 15second interface material. Such acid functionality may be introduced as part of the second interface material formulation or introduced later, after curing, using an acid water solution. For example, the first step may be to put the threedimensional object with its support in a water solution containing a SBC, then to place the same object in an acidic solution. The acid will start to decompose the SBC and produces gas (bubbles). In another embodiment, the substance that liberates gas is already present in the formulation of the second interface material. For example, the second interface material may contain calcium carbonate as a solid filler. In that case, the trigger is the introduction of the second interface material in a water or acidic solution. 30 It should be clear that a BRS is not limited to a sodium bicarbonate or calcium carbonate and an acidic water solution. Other chemical reagents and reactions may be used to achieve the same result—the production of bubbles inside the matrix of the second interface material. For example, the SBC may be any alkaline metal or alkaline earth metal carbonate or bicarbonate.

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In one embodiment of the present invention, the composition further includes, inter alia, a filler. In another embodiment, the filler includes particles having a diameter of less than 1 micron.

In one embodiment of the present invention, the composition further includes a low molecular weight polymer. In one embodiment of the present invention, the first viscosity composition suitable for support in building a threedimensional object is about 30-200 cps.

In one embodiment of the present invention, the second viscosity of the composition suitable for support in building a three-dimensional object is lower than 20 cps. In another embodiment, the second viscosity is between 10 and 17 cps. In another embodiment, the second viscosity is between 12 and 16 cps. Having these viscosities, the first and second interface material may be distinguished from certain prior art formulations designed for ink-jet printing, which may have low viscosity at room temperature, the temperature at which the printing is typically conducted. High viscosity at room temperature may be a desirable property for three-dimensional objects, a feature that may be lacking in the prior art formulations. In one embodiment of the present invention, the composition further includes, inter alia, a component able to produce gas upon exposure to water or to an alkaline or acidic water solution. In another embodiment, the component is sodium bicarbonate, calcium bicarbonate or a combination thereof. Other suitable substances may be used. In one embodiment of the present invention, the second interface composition further includes, inter alia, a pigment, a dye or a combination thereof. In another embodiment, the pigment is a white pigment, an organic pigment, an inorganic pigment, a metal pigment or a combination thereof.

Examples of formulations of the second interface material are provided hereinbelow in Table 5 and Table 6, to which reference is now made. Tables 5 and 6 display various formulations that are suitable for use as the second interface material. The individual substances, suppliers, combinations, etc., are given by way of example only.

In one embodiment of the present invention, the non-curable component is a non-water miscible compound. In another embodiment, the non-water miscible compound is paraffin oil.

TABLE 5

|   | Examples of Characteristic Formulation Components of Second Interface Material |                                          |                                             |                                      |  |  |  |  |  |
|---|--------------------------------------------------------------------------------|------------------------------------------|---------------------------------------------|--------------------------------------|--|--|--|--|--|
| # | Trade Name                                                                     | Chemical Type                            | Function in the formulation                 | Supplier                             |  |  |  |  |  |
| A | SR-610                                                                         | Polyethylene Glycole (600)<br>Diacrylate | Oligomer                                    | Sartomer                             |  |  |  |  |  |
| В | Bisomer PEA6                                                                   | Polyethylene Glycole<br>monoacrylate     | Water swelling/<br>sensitive Oligomer       | Laport                               |  |  |  |  |  |
| С | PEG 400                                                                        | Polyethylene Glycole 400                 | Polymer<br>(hydrophilic and<br>plasticizer) | Aldrich                              |  |  |  |  |  |
| D | Irgacure 907                                                                   | alpha-Amino Ketone                       | Free radical photo-initiator                | Ciba<br>Specialties<br>Chemical Inc. |  |  |  |  |  |
| Е | BP                                                                             | Benzophenone                             | Type I<br>Free radical<br>photo-initiator   | Satomer                              |  |  |  |  |  |

|   |            |                              | Type II            |            |
|---|------------|------------------------------|--------------------|------------|
| F | Triethanol | Ternary Amine                | Free radical       | Aldrich    |
|   | Amine      |                              | Coinitiator for    |            |
|   |            |                              | Type II photo-     |            |
|   |            |                              | initiator          |            |
| G | Byk 307    | Silicone Surface Additive    | Surface agent      | Byk Chemie |
| Η | MEHQ       | 4-Methoxy phenol             | Inhibitor          | Sigma      |
|   |            |                              | (thermal           |            |
|   |            |                              | stabilizer)        |            |
| Ι | PEG UA     | Polyethylene glycol urethane | Water swelling/    | Home made  |
|   |            | diacrylate                   | sensitive oligomer |            |
|   |            | -                            | -                  |            |

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TABLE 5-continued

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#### Examples of Characteristic Formulation Components of Second Interface Material

| # | Trade Name          | Chemical Type                            | Function in the formulation           | Supplier     |
|---|---------------------|------------------------------------------|---------------------------------------|--------------|
| J | AP                  | Partially acrylated polyol               | Water swelling/<br>sensitive oligomer | Home made    |
| Κ | Betha-CEA           | Betha-caboxyethyl acrylate               | Acidic monomer                        |              |
| М | CHVE                | 1,4-Cyclohexane dimethanol divinyl ether | Vinyl ether<br>monomer                | ISP          |
| Ν | Tone polyol<br>0301 | Caprolactone polyol                      | Polyol<br>(plasticizer)               | Union Cabide |
| Ο | Paraffin oil        | Paraffin oil                             | plasticizer                           | Oldrich      |
| Р | methoxycarbo        | methoxypolyethylene glycol               | Polymer<br>(hydrorphilic and          |              |

|   | wax 500 and |                   | (inveroprine and |             |
|---|-------------|-------------------|------------------|-------------|
|   | 1000        |                   | plasticizer)     |             |
| Q | SR 506      | Isoborny Acrylate | monomer          | Cray Valley |
|   |             |                   |                  |             |

#### TABLE 6

Examples of Possible Formulation Compositions of Second Interface Material

Example A B C D E F G H I J K L M O Q N

| 1<br>2<br>3 | X<br>X | Х | X<br>X<br>X | X<br>X | Х  | Х  | X<br>X<br>X | X<br>X<br>X |   |   |   |   |   |   |   |   |
|-------------|--------|---|-------------|--------|----|----|-------------|-------------|---|---|---|---|---|---|---|---|
|             |        |   |             | Λ      | 37 | 37 |             |             |   |   |   |   |   |   |   |   |
| 4           |        | Х | Х           |        | X  | Х  | Х           | Х           |   |   |   |   |   |   |   |   |
| 5           | Х      | Х | Х           | Х      |    |    | Х           | Х           |   |   |   |   |   |   |   |   |
| 6           | Х      | Х | Х           |        | Х  | Х  | Х           | Х           |   |   |   |   |   |   |   |   |
| 7           |        |   | Х           |        |    |    | Х           | Х           | Х |   |   |   |   |   |   |   |
| 8           |        |   | Х           | Х      |    |    | Х           | Х           |   | Х |   |   |   |   |   |   |
| 9           |        | Х | Х           | Х      |    |    | Х           | Х           |   |   | Х |   |   |   |   |   |
| 10          |        |   | Х           | Х      |    |    | Х           | Х           |   |   |   | Х |   |   |   |   |
| 11          |        |   | Х           | Х      |    |    | Х           | Х           |   |   |   |   |   |   |   |   |
| 12          | Х      |   |             | Х      |    |    | Х           | Х           |   |   |   | Х | Х |   |   |   |
| 13          |        | Х |             | Х      |    |    | Х           | Х           |   | Х |   |   |   |   |   | Х |
| 14          |        | Х | Х           | Х      | Х  | Х  | Х           | Х           |   | Х | Х |   |   |   |   | Х |
| 15          |        |   |             | Х      |    |    |             |             |   |   |   |   |   | Х | Х |   |

A formulation of the second interface material is presented in entry No. 3 of Table 6. According to this embodiment of the present invention, the second interface material includes: a water swelling oligomer, which may be any water swelling oligomer as defined hereinabove, and which may be polyethylene glycol; a non-curable component, which may be any non-curable component as defined hereinabove, and which may be polyethylene glycole; a radical photo-initiator, which may be any radical photo-initiator as defined hereinabove, and which may be alpha-amino ketone; a surface agent, which may be a silicone surface additive; and an inhibitor, which may be 4-methoxyphenol.

Another formulation of the second interface material is presented in entry No. 4 of Table 6. According to this embodiment of the present invention, the second interface material includes: a water swelling oligomer, which may be any water swelling oligomer as defined hereinabove, and which may be 55 polyethylene glycole monoacrylate; a non-curable component, which may be any non-curable component as defined hereinabove, and which may be polyethylene glycole; a radical photo-initiator, which may be any radical photo-initiator as defined hereinabove, and which may be benzophenone; a 60 co-initiator, which may be any co-initiator as defined hereinabove, and which may be triethanolamine; a surface agent, which may be a silicone surface additive; and an inhibitor, which may be 4-methoxyphenol. The first interface material and the second interface mate- 65 rial are suitable for use in, for example, the method for threedimensional printing which is described in U.S. patent appli-

cation Ser. No. 09/412,618, assigned to the Assignees of the present application and is incorporated herein by reference. Other methods may be used.

Briefly, the method according to one embodiment includes: dispensing a first interface material from a printing head; dispensing a second interface material from the printing head; and combining the first interface material and the second interface material in pre-determined proportions to a produce a multiplicity of construction layers for forming the three-dimensional model.

The method (FIG. 3) according to an embodiment of the 50 present invention includes dispensing a first composition suitable for building a three-dimensional object from a dispenser (102), dispensing a second composition suitable for support in building a three-dimensional object from a dispenser (104), combining the first composition and the second composition in pre-determined proportions to produce a multiplicity of construction layers for forming the three-dimensional object (106), curing the first composition resulting in a solid form (108), and irradiating or curing second composition resulting in a liquid, a solid or a semi-solid form (110). Of course, the method may include other steps or series of steps. One embodiment of the present invention further provides a method for the preparation of a three-dimensional object by three-dimensional printing, the method may include the steps of dispensing a first composition suitable for building a threedimensional object from a dispenser, the first composition may include, inter alia, a curable component, having a func-

tional group, wherein if the functional group is a polymeriz-

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able reactive functional group, then the functional group is a (meth)acrylic functional group, a photo-initiator, a surfaceactive agent; and a stabilizer, dispensing a second composition suitable for support in building a three-dimensional object from a dispenser, the second composition may include a non-curable component, a curable component, wherein the non-curable component is not reactive with the curable component, a surface-active agent and a stabilizer, combining the first composition and the second composition in pre-determined proportions to produce a multiplicity of construction <sup>10</sup> layers for forming the three-dimensional object, whereby the first composition is cured resulting in a solid form, and whereby the second composition is irradiated or cured resulting in a liquid, a solid or a semi-solid form. In one embodiment of the present invention, the method may further include the step of generating data for a predetermined combination of the first composition and the second composition to produce a multiplicity of support layers for supporting the three-dimensional object. 20 In one embodiment of the present invention, the method may further include the step of generating data for a predetermined combination of the first composition and the second composition to produce a multiplicity of release layers for releasing the three-dimensional object from the support 25 layers. In one embodiment of the present invention, the first composition and the second composition are dispensed simultaneously. In another embodiment, the first composition and the second composition are dispensed sequentially. In another 30 embodiment, the first composition is dispensed first. In another embodiment, the second composition is dispensed first. In another embodiment, more than one first composition is used. In another embodiment, the more than one second composition is used. In accordance with one embodiment of the present invention, the method further includes the step of curing the first interface material. Further, when the second interface material includes a curable component, the method may further include the step of curing the second interface material. Cur- 40 ing may be carried out for example, as described in U.S. Pat. No. 6,658,314. For example, the curing method is by radiation, such as Ultraviolet (UV) and/or Visible (Vis) and/or Infra Red (IR) and/or UV-Vis radiation and/or Electron Beam (EB). In one embodiment of the present invention, the curing 45 method is UV-Vis radiation. Other suitable curing methods may be used. In operation, in order to obtain layers of different modulus of elasticity and a different strength, the first interface material and the second interface material are combined in pre- 50 determined proportions. For example, in order to obtain layers having a higher modulus of elasticity and a higher strength such as the construction layers, a suitable combination that contains mostly the first interface material may be used. Further, in order to obtain layers having a lower modulus of 55 elasticity and a lower strength such as the release layers, a suitable combination that includes mostly the second interface material may be used. By way of example, in order to produce the construction layers and/or the support layers, a combination that includes 60 90-100% of the first interface material and 0-10% of the second interface material may be used. Further, in order to produce the release layers, a combination that includes 0-10% of the first interface material and 90-100% of the second interface material may be used. In another embodiment, in 65 order to produce support layers that have a lower modulus of elasticity and a lower strength than the construction layers, a

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combination that includes 30-70% of the first interface material and 70-30% of the second interface material may be used.

Thus a three-dimensional object is produced which is included of a core consisting of a multiplicity of construction layers. The construction layers are formed by combining predetermined proportions of the first interface material and the second interface material.

One embodiment of the present invention further provides a three-dimensional object comprised of a multiplicity of construction layers, wherein the construction layers are prepared by combining pre-determined proportions of a first composition and a second composition according to the invention. In another embodiment the three-dimensional object is comprised of a core consisting of a multiplicity of construction layers, wherein the construction layers are prepared by combining pre-determined proportions of a first composition and a second composition according to the invention.

One embodiment of the present invention provides a threedimensional object including the composition according the invention.

In one embodiment of the present invention, the threedimensional object further includes a multiplicity of support layers for supporting the core. The support layers are prepared by combining pre-determined proportions of the first interface material and the second interface material. The support layers may be designed exactly like to construction layers, or may be designed to be weaker (lower modulus of elasticity) than the construction layers.

In one embodiment of the present invention, the threedimensional object may further include a multiplicity of support layers for supporting the core, wherein the support layers are prepared by combining pre-determined proportions of the first composition and the second composition. In another

embodiment, the support layers support the construction layers.

In another embodiment, the support layers have the same strength as the construction layers. In another embodiment, the support layers have the same modulus of elasticity as the construction layers. In another embodiment, the support layers have a lower modulus of elasticity and/or a lower strength than the construction layers.

In one embodiment of the present invention, the threedimensional object further includes a multiplicity of release layers for releasing the support layers from the construction layers. In one embodiment of the present invention, the release layers are positioned between the support layers and the construction layers. The release layers are prepared by combining pre-determined proportions of the first interface material and the second interface material.

In one embodiment of the present invention, the threedimensional object may further include a multiplicity of release layers for releasing the support layers from the core, wherein the release layers are positioned between the support layers and the construction layers; wherein the release layers are prepared by combining pre-determined proportions of the first composition and the second composition. In another embodiment, the release layers have a lower modulus of elasticity and/or a lower strength than the construction layers and the support layers. It will be appreciated by persons skilled in the art that the present invention is not limited by what has been particularly shown and described herein above and that numerous modifications, all of which fall within the scope of the present invention, exist. Rather, the scope of the invention is defined by the claims which follow:

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What is claimed is:

1. A radiation curing composition suitable for building a three dimensional object by a solid freeform method, the composition comprising:

one or more meth(acrylic)mono-functional monomers <sup>5</sup> where a respective polymer has a Glass Transition Temperature higher than about 60° C.;

one or more meth(acrylic)di-functional oligomers where a respective polymer has a Glass Transition Temperature lower than about 40° C.; and

an acrylated silicon oligomer,

wherein the concentration of said one or more meth (acrylic)mono-functional monomers is at least 20% by

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**5**. The composition of claim **1**, wherein the composition has a viscosity lower than 20 cps at a temperature higher than the ambient temperature and after curing, the composition results in a solid form.

6. The composition of claim 1, further comprising a molecule having one or more epoxy substituents, a molecule having one or more vinyl ether substituents, vinylcaprolactam, vinylpyrolidone, or any combination thereof.

7. The composition of claim 1, further comprising a photo-10 initiator.

**8**. The composition of claim **1**, further comprising a sulfurcontaining component.

**9**. The composition of claim **8**, wherein said sulfur-containing component is beta mercaptopropionate, mercaptoacto etate, alkane thiols or any combination thereof.

weight relative to the total weight of the composition and the concentration of said one or more meth(acrylic)difunctional oligomers is at least 20% by weight relative to the total weight of the composition; and wherein the composition has a viscosity of about

50-500cps at ambient temperature.

2. The composition of claim 1, wherein the amount of the silicon oligomer is about 5% by weight relative to the total weight of the composition.

**3**. The composition of claim **1**, wherein upon curing the composition results in a material having improved impact strength relative to a similar composition without the acrylated silicon oligomer.

**4**. The composition of claim **1**, wherein said the acrylated silicon oligomer promotes phase separation during curing of said composition.

10. The composition of claim 1, further comprising at least one pigment and at least one dispersant.

**11**. The composition of claim **1**, further comprising a dye.

12. The composition according to claim 1, wherein the Glass Transition Temperature of the mono-functional monomer is higher than  $70^{\circ}$  C.

**13**. The composition of claim **1**, wherein the viscosity at ambient temperature is above 80 cps.

14. The composition of claim 1, wherein viscosity at ambi-25 ent temperature is about 300 cps.

15. The composition of claim 1, wherein the di-functional oligomer is urethane-acrylate oligomer or urethane meth-acrylate oligomer.

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